Silicas precipitated by a specific method for rubber applications

The present invention relates to precipitated silicas which have extremely low microporosity and high rubber activity, to a process for their preparation, and to their use as filler for rubber mixtures.

The use of precipitated silicas in elastomer mixtures, such as tire tread mixtures, has been known for a long 10 time (EP 0 501 227). There are stringent requirements relating to the use of silicas as a reinforcing filler in rubber mixtures, for example those used to produce pneumatic tires and technical rubber products. They have to have low density and have good capability for incorporation and dispersion in the rubber, and undergo chemical bonding to the rubber in association with a reagent, preferably bifunctional a coupling organosilicon compound, this reaction leading to the desired high level of reinforcement of the rubber 20 reinforcement property of The mixture. particular be associated with high static stress values and a low abrasion value. The particle size, surface morphology, surface activity, and also the binding power of the coupling reagent, are of particularly 25 decisive importance in relation to the reinforcing property of the silicas.

Precipitated silicas with primary particle diameters of in rubber are usually used to 50 nm from 10 30 One way of determining the applications. particle diameter is by means of transmission electron evaluation (R. H. Lange, image spectroscopy (TEM) J. Bloedorn: "Das Elektronenmikroskop, TEM + REM" [The Thieme Verlag, electron microscope, TEM + SEM] 35 Stuttgart, New York (1981)). The surface areas measured via the adsorption of the small nitrogen molecule N_2 , and are termed BET surface area below

(S. Brunauer, P. H. Emmett, E. Teller, "Adsorption of Gases in Multimolecular Layers", J. Am. Chem. Soc. 60, 309 (1938)), and are typically in the range from 50 to 200 m^2/g , preferably from 110 to 180 m^2/g . The surface area accessible to the rubber, hereinafter termed the 5 surface area, is determined by means of the larger N-cetyl-N,N,N-trimethylthe adsorption of ammonium bromide molecule (See Janzen, Kraus, Rubber Chem. Technol. 44, 1287 (1971)). Detailed information describing exactly how the measurements are made is 10 given in the following text. The difference between the BET surface area and the CTAB surface area can therefore be termed the microporosity, and represents the silica surface area inaccessible to the rubber and for reinforcement. ineffective 15 therefore commercially available active silicas the BET/CTAB ratio is usually > 1.05, indicating a correspondingly high degree of microporosity.

In addition, the skilled worker is aware that low-molecular-weight compounds, e.g. the bifunctional organosilicon compounds and vulcanization accelerators, can undergo physi- and chemisorption within the pores of the microporous silica, and thus lose their ability to carry out their function as coupling agent for the rubber or vulcanization accelerator for crosslinking of the rubber.

The skilled worker is also aware that the coupling reagent, usually a bifunctional organosilicon compound known from S. Wolff, "Chemical Aspects of Rubber Reinforcement by Fillers", Rubber Chem. Technol. 69, 325 (1996), is intended to modify the rubber-active surface quantitatively and with maximum uniformity. The modification may take place via precoating of the silica in bulk or in solution/suspension (ex situ) (U. Görl, R. Panenka, "Silanisierte Kieselsäuren - Eine neue Produktklasse für zeitgemässe Mischungsent-

wicklung" (Silanized silicas - a new product class for advanced compound development), Kautsch. Gummi Kunstst. 46, 538 (1993)), or else during the mixing process (in situ) (H.-D. Luginsland, "Processing of Silica/Silane-Filled Tread Compounds", paper No. 34 presented at the ACS Meeting, April 4-6, 2000, Dallas, Texas/USA), in situ modification being the preferred and usually used process. To ensure rapid and quantitative silanization of the rubber-active surface, it is therefore necessary to enrich the surface in a controlled manner with 10 reactive silanol groups to which the organosilicon The interaggregate structure of compound can bond. silicas is usually determined by means of the DBP test (J. Behr, G. Schramm, "Über die Bestimmung der Ölzahl Kautschukfüllstoffen mit . dem Brabender-15 Plastographen" (Determination of the oil value of rubber fillers using the Brabender Plastograph], Gummi Asbest Kunstst. 19, 912 (1966)). A high DBP value is needed to ensure ideal dispersion of the filler in the rubber. A moisture level > 4% is needed to ensure rapid 2.0 and complete silanization of the silica surface with an organosilicon compound (U. Görl, A. Hunsche, A. Müller, H. G. Koban, "Investigations into the Silica/Silane Technol. 70, Reaction System", Rubber Chem. (1997)). 25

JP 2002179844 discloses rubber mixtures with silicas in which the difference between BET surface area and CTAB surface area of the silicas is smaller than 15 m²/g. No mention is made of the high Sears value V2, this being a measure of the number of silanol groups needed for high activity with respect to the bifunctional organosilicon compound, or of the high DBP value needed for very good dispersion in the rubber. However, these two values, alongside the low microporosity, are decisive for a high degree of reinforcement of the rubber mixture and therefore good road abrasion properties.

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Moisture level

The skilled is worker is aware of various processes for preparing silicas. For example, EP 0 937 755 describes precipitations at constant pH. DE 101 24 298 discloses silicas precipitated at constant cation excess. DE 101 12 441 A1, EP 0 755 899 A2, and US 4 001 379 describe precipitations at constant alkali value (AV value).

Silicas for rubber applications are generally prepared by a process in which the precipitation takes place at temperatures of from 60 to 95°C and at a non-constant pH of from 7 to 10, see EP 0 901 986 A1, for example.

It is an object of the present invention to provide novel, readily dispersible precipitated silicas which can be incorporated in elastomer mixtures and which improve their properties. A further intention is to provide a process for preparing these silicas.

The present invention therefore provides readily dispersible precipitated silicas characterized by:

20	dispersible precipitated sil	icas characterized by.
	CTAB surface area	$100-200 \text{ m}^2/\text{g}$, the preferred
	•	ranges being $100-160 \text{ m}^2/\text{g}$,
		$100-130 \text{ m}^2/\text{g}, \text{ and } 110-125$
		m^2/g ,
25	BET/CTAB ratio	0.8-1.05, preferred ranges
		0.90-1.03, 0.93-1.03, and
		particularly 0.96-1.03,
	DBP value	210-280 g/(100 g), prefer-
		red range 250-280 g/
30		(100 g),
	Sears value V ₂	10-30 m1/(5 g), the
		preferred ranges being
		20-30 ml/(5 g)and 21 to 28
		(ml/5 g),

The precipitated silicas of the invention may moreover preferably have one or more of the following physico-

4-8%.

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chemical parameters:

80-210 m²/a, BET surface area precipitated silicas in a first embodiment having a 5 $80-110 \text{ m}^2/\text{g}$ and BET of second in a those embodiment having a BET of $110-150 m^2/g$, preferably $110-129 \text{ m}^2/\text{g}$ 10 Ratio of Sears value V₂ $0.140-0.370 \text{ ml/}(5 \text{ m}^2)$, the to BET surface area ranges being preferred $0.150-0.320 \text{ ml/}(5 \text{ m}^2)$, $0.170-0.320 \text{ ml/}(5 \text{ m}^2)$, 15 $m1/(5 m^2)$, 0.180-0.300 particularly preferably $0.190-0.280 \text{ ml/}(5 \text{ m}^2) \text{ and}$ $0.190-0.240 \text{ ml/}(5 \text{ m}^2)$,

20 Primary particle diameter 10-50 nm.

In one preferred embodiment, the precipitated silicas of the invention have a Sears value V_2 of 22-25 ml/(5 g), the Sears value V_2 in another embodiment being 25-28 ml/(5 g).

The precipitated silicas of the invention have not only a high absolute number of silanol groups (Sears value V_2), but also, when comparison is made with prior-art precipitated silicas, a markedly increased ratio of the Sears value V_2 to the BET surface area. This means that the precipitated silicas of the invention in particular have a very high number of silanol groups based on the total surface area.

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The precipitated silicas of the invention have not only an increased number of silanol groups but also low microporosity, i.e. a very low ratio of BET to CTAB.

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The combination of the features mentioned, in particular the high ratio of Sears value V_2 to BET, gives the precipitated silicas of the invention excellent suitability as reinforcing fillers for elastomers. These precipitated silicas of the invention have increased rubber activity, and exhibit very good dispersion behavior and a low vulcanization time.

The present invention further provides a process which can be used to prepare the precipitated silicas of the invention with

CTAB surface area $100-200 \text{ m}^2/\text{g}, \text{ the preferred}$ $\text{ranges being } 100-160 \text{ m}^2/\text{g},$ $100-130 \text{ m}^2/\text{g}, \text{ and}$ $110-125 \text{ m}^2/\text{g},$ BET/CTAB ratio 0.8-1.05, preferred ranges 0.90-1.03, 0.93-1.03, and

particularly 0.96-1.03,

DBP value 210-280 g/(100 g),

preferred range 250-280 g/(100 g),

25 . 28 (m1/5 g), Moisture level 4-8%,

and, where appropriate, with one or more of the following physico-chemical parameters

30 BET surface area $80-210 \quad m^2/g, \quad \text{the} \\ \text{precipitated silicas in a} \\ \text{first embodiment having a} \\ \text{BET of } 80-110 \quad m^2/g \quad \text{and} \\ \text{those in a second} \\ \text{embodiment having a BET of} \\ 110-150 \quad m^2/g, \quad \text{preferably} \\ \end{cases}$

 $110-129 \text{ m}^2/\text{g}$

Ratio of Sears value V2

to BET surface area

0.140-0.370 ml/(5 m²), the preferred ranges being 0.150-0.320 ml/(5 m²), 0.170-0.320 ml/(5 m²).

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 $\begin{array}{lll} 0.170 - 0.320 & ml/(5 & m^2) \,, \\ 0.180 - 0.300 & ml/(5 & m^2) \,, \\ particularly & preferably \\ 0.190 - 0.280 & ml/(5 & m^2) & and \\ 0.190 - 0.240 & ml/(5 & m^2) \,, \end{array}$

primary particle diameter which comprises first

a) taking an aqueous solution of an alkali metal silicate or alkaline earth metal silicate and/or of an organic and/or inorganic base with pH from 7 to 14 as initial charge,

10-50 nm,

- b) metering water glass and an acidifier simultaneously into this initial charge at from 55 to 95°C, with stirring, for from 10 to 120 minutes, preferably from 10 to 60 minutes,
- 20 g) continuing to stir the resultant suspension at from 80 to 98°C, for from 1 to 120 minutes, preferably from 10 to 60 minutes,
 - h) using an acidifier to acidify to pH of from 2.5 to 5, and
- 25 i) filtering, washing, drying and, where appropriate, pelletizing.

The initial charge may amount to about 20, 30, 40, 50, 60, 70, 80, or 90% of the final volume for the precipitation. The basic compounds for the initial 30 charge are in particular selected from the group of the earth alkaline hydroxides, metal alkali metal carbonates, alkali metal alkali hydroxides, silicates. and alkali metal hydrogencarbonates, Preference is given to the use of water glass and/or 35 sodium hydroxide solution. The pH of the initial charge is from 7 to 14, preferably from 10 to 11.

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The manner of adding the water glass and the acidifier during step b) may be such that the pH of the reaction solution is held constant or remains variable. However, it is also possible for the precipitation to take place at a constant alkali value. If the precipitation takes place at a constant pH, this is preferably in the range from 7 to 11. Precipitations at a constant alkali value preferably take place at an alkali value of from 15 to 40.

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After step b) the process of the invention may comprise one or two hold points, viz.:

- c) stopping the feed for from 30 to 90 minutes while maintaining the temperature, and
- 15 d) then, where appropriate, simultaneously feeding water glass and an acidifier at the same temperature, with stirring, for from 20 to 120 minutes, preferably from 20 to 80 minutes.
- 20 Following step b) or d) of the process of the invention, addition of water glass is stopped and, if the desired pH has not yet been achieved, in step
- e) the acidifier feed is continued at an identical or different rate until a pH of 5 to 11, preferably from 7 to 10, has been achieved.

In order to vary the concentration of the water glass available for the reaction, where appropriate,

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f) the pH may be adjusted to from 3 to 11 by adding further acidifier, and/or raised again to values of from 8 to 14, preferably 9-10, and particularly 7-11, via feed of one or more basic compounds from the group of the alkali metal hydroxides, alkaline earth metal hydroxides, alkali metal carbonates, alkali metal hydrogencarbonates, and alkali metal silicates. Use of water glass and/or sodium

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hydroxide solution is preferred.

The feed of the acidifier in step e) or of the basic compound in step f) may take place at from 40 to 98°C. The feed preferably takes place at from 55 to 95°C and particularly preferably at the temperature also selected for step b) or d).

Another hold point may optionally be incorporated 10 between steps d) and e).

Another addition of organic or inorganic salts may moreover optionally be incorporated during one or more of steps a) to h). This addition may be carried out in solution or in solid form, in each case continuously over the period of addition of the water glass and of the acidifier, or may be batch addition. It is also possible for the salts to be dissolved in one or both of the components and then to be added simultaneously with these.

The inorganic salts preferably used are alkali metal salts or alkaline earth metal salts. In particular, use may be made of any of the combinations of the following ions:

 $\begin{array}{l} \text{Li}^{+}, \ \text{Na}^{+}, \ \text{K}^{+}, \ \text{Rb}^{+}, \ \text{Be}^{2+}, \ \text{Mg}^{2+}, \ \text{Ca}^{2+}, \ \text{Sr}^{2+}, \ \text{Ba}^{2+}, \ \text{H}^{+}, \ \text{F}^{-}, \ \text{Cl}^{-}, \\ \text{Br}^{-}, \ \text{I}^{-}, \ \text{SO}_{3}^{2-}, \ \text{SO}_{4}^{2-}, \ \text{HSO}_{4}^{-}, \ \text{PO}_{3}^{3-}, \ \text{PO}_{4}^{3-}, \ \text{NO}_{3}^{-}, \ \text{NO}_{2}^{-}, \ \text{CO}_{3}^{2-}, \\ \text{HCO}_{3}^{-}, \ \text{OH}^{-}, \ \text{TiO}_{3}^{2-}, \ \text{ZrO}_{3}^{2-}, \ \text{ZrO}_{4}^{4-}, \ \text{AlO}_{2}^{-}, \ \text{Al}_{2}\text{O}_{4}^{2-}, \ \text{BO}_{4}^{3-}. \end{array}$

Suitable organic salts are the salts of formic, acetic, or propionic acid. Cations which may be mentioned are the abovementioned alkali metal ions or alkaline earth metal ions. The concentration of these salts in the solution added may be from 0.01 to 5 mol/l. The inorganic salt preferably used is Na₂SO₄.

The manner of introducing the acidifier in steps b) and d) may be identical or different, i.e. its concentration and/or feed rate may be identical or

different.

Similarly, a manner of introducing the water glass to the reaction in steps b) and d) may be identical or different.

In one particular embodiment, the manner of introducing the components acidifier and water glass in steps b) and d) is such that the feed rate in step d) is from 125 to 140% of the feed rate in step b), each of the components being used in equimolar concentration in the two steps. It is preferable for the components to be added with identical concentration and feed rate.

Besides water glass (sodium silicate solution), use may also be made of other silicates, such as potassium silicate or calcium silicate. Acidifiers which may be used, besides sulfuric acid, are HCl, HNO₃, H₃PO₄, or CO₂.

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The filtration and long-term or short-term drying of the silicas of the invention are matters familiar to the skilled worker and can be found in the documents mentioned, for example.

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The silica precipitated is preferably dried in a pneumatic drier, spray drier, disk drier, belt drier, rotating-tube drier, flash drier, spin flash drier, or spray tower. These drying methods include operation using an atomizer, using a single- or twin-fluid nozzle, or using an integrated fluidized bed. Where appropriate, after the drying process grinding may take place, and/or a roller compactor may be used for palletizing. After the drying step, the grinding, or the pelletization, the precipitated silica of the invention preferably has a particle shape with an average diameter above 15 µm, in particular above 80 µm, particularly preferably above 200 µm (determined

ISO 2591-1, December 1988). The precipitated silicas of the invention particularly preferably take the form of a powder with an average diameter above 15 $\mu \rm m$, or take the form of substantively round particles with an average diameter above 80 $\mu \rm m$ (microbeads), or take the form of pellets with an average diameter \geq 1 mm.

The present invention also provides the use of the precipitated silicas of the invention, with a

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	CTAB surface area	$100-200 \text{ m}^2/\text{g}$, the preferred
		ranges being $100-160 \text{ m}^2/\text{g}$,
		$100-130$ m^2/g , and
		$110-125 \text{ m}^2/\text{g},$
15	BET/CTAB ratio	0.8-1.05, preferred ranges
		0.90-1.03, 0.93-1.03, and
		particularly preferably
		0.96-1.03,
	DBP value	210-280 g/(100 g),
20		preferred range
		250-280 g/(100 g),
	Sears value V ₂	10-30 ml/(5 g), the
		preferred ranges being
		20-30 ml/(5 g)and 21 to 28
25		(m1/5 g),
_•	Moisture level	4-8%

and also, where appropriate, with one or more of the following physico-chemical parameters

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	BET surface area	80-210	m^2/g ,	the
		precipita	ted silic	as in a
		first em	bodiment 1	naving a
		BET of	80-110 n	n ² /g and
35		those	in a	second
	·	embodimen	t having	a BET of
	•	110-150	m^2/g , pr	referably
		110-129 m	ι²/g,	

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Ratio of Sears value V_2 to BET surface area

10 primary particle diameter 10-50 nm

for preparing elastomer mixtures, vulcanizable rubber mixtures, and/or other vulcanisates.

The invention therefore also provides elastomer mixtures, vulcanizable rubber mixtures, and/or other vulcanizates which comprise the silica of the invention, examples being moldings, such as pneumatic tires, tire treads, cable sheathing, hoses, drive belts, conveyor belts, roller coverings, tires, shoe soles, ring seals, and damping elements.

The silicas of the invention may also be used in any of the application sectors which usually use silicas, e.g. in battery separators, as antiblocking agent, 25 matting agent in inks and paints, as carrier for agricultural products and for feeds, in coatings, powders, in fire-extinguisher inks, printing plastics, in the non-impact printing sector, in paper pulp, or in the personal care sector, or in specialty 30 applications. Use in the non-impact printing sector, for example in inkjet printing, is use of the silicas of the invention in

- printing inks for thickening or for preventing
 splashing and offset,
 - paper, as filler or coating pigment, blueprint paper, heat-sensitive paper, thermal sublimation for preventing strike-through of printing inks, for

improvement in contrast and image background uniformity, and for improvement in dot definition and color brilliance.

Use in the personal care sector is use of the silicas of the invention as filler or thickener, e.g. in the pharmaceutical sector or the body-care sector.

The silica of the invention may optionally be modified with silanes or organosilanes of the formulae I to III

10 $[SiR^{1}_{n}(OR)_{r}(Alk)_{m}(Ar)_{p}]_{q}[B]$ (I), $SiR^{1}_{n}(OR)_{3-n}(Alkyl)$ (II),

or $SiR^{1}_{n}(OR)_{3-n}(Alkenyl)$

(III),

15 where

B is -SCN, -SH, -Cl, $-NH_2$, $-OC(O)CHCH_2$, $-OC(O)C(CH_3)CH_2$ (if q=1), or $-S_w$ - (if q=2), B being chemically bonded to Alk,

R and R¹ are an aliphatic, olefinic, aromatic, or arylaromatic radical having 2-30 carbon atoms, optionally with substitution by the following groups: the hydroxyl, amino, alcoholate, cyanide, thiocyanide, halo, sulfonic acid, sulfonic ester, thiol, benzoic acid, benzoic ester, carboxylic acid, carboxylic ester, acrylate, methacrylate, or organosilane radical, where the meaning or substitution of R and R¹ may be identical or different,

n is 0, 1, or 2,

30 Alk is a bivalent unbranched or branched hydrocarbon radical having from 1 to 6 carbon atoms,

m is 0 or 1,

Ar is an aryl radical having from 6 to 12 carbon atoms, preferably 6 carbon atoms, which may have substitution by the following groups: the hydroxyl, amino, alcoholate, cyanide, thiocyanide, halo, sulfonic acid, sulfonic ester,

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thiol, benzoic acid, benzoic ester, carboxylic acid, carboxylic ester, or organosilane radical,

p is 0 or 1; with the proviso that p and n are not simultaneously 0,

q is 1 or 2,

w is a number from 2 to 8,

r is 1, 2, or 3, with the proviso that r + n + m + p = 4,

10 Alkyl is a monovalent unbranched or branched saturated hydrocarbon radical having from 1 to 20 carbon atoms, preferably from 2 to 8 carbon atoms,

Alkenyl is a monovalent unbranched or branched unsaturated hydrocarbon radical having from 2 to 20 carbon atoms, preferably from 2 to 8 carbon atoms.

The silica of the invention may also be modified with organosilicon compounds of the composition $\text{SiR}^2_{4-n}X_n$ 20 (where n = 1, 2, 3, 4), $[SiR^2_xX_yO]_z$ (where $0 \le x \le 2$; $0 \le x \le 2$) $y \le 2$; $3 \le z \le 10$, where x + y = 2), $[SiR^2_xX_yN]_z$ (where 0 $\leq x \leq 2$; $0 \leq y \leq 2$; $3 \leq z \leq 10$, where x + y = 2), $SiR^{2}_{n}X_{m}OSiR^{2}_{o}X_{p}$ (where $0 \le n \le 3$; $0 \le m \le 3$; $0 \le o \le 3$; 0 \leq p \leq 3, where n + m = 3, o + p = 3), $SiR^{2}{}_{n}X_{m}NSiR^{2}{}_{o}X_{p}$ 25 (where $0 \le n \le 3$; $0 \le m \le 3$; $0 \le o \le 3$; $0 \le p \le 3$, where n + m = 3, o + p = 3), $SiR^2_n X_m [SiR^2_x X_y O]_z SiR^2_o X_p$ (where $0 \le 1$) $n \le 3$; $0 \le m \le 3$; $0 \le x \le 2$; $0 \le y \le 2$; $0 \le o \le 3$; $0 \le p$ \leq 3; 1 \leq z \leq 10000, where n + m = 3, x + y = 2, o + p = 3). These compounds may be linear, cyclic, or branched, 30 silane, silazane, or siloxane compounds. R² may be substituted and/or unsubstituted alkyl and/or aryl radicals, in each case having 1-20 carbon atoms, which may have substitution by functional groups, for example by the hydroxyl group, by the amino group, 35 polyethers, such as ethylene oxide and/or propylene oxide, or by halide groups, such as fluoride. R^2 may also include groups such as alkoxy, alkenyl, alkynyl,

preferably being methyl.

preferably being methyl.

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and aryl groups, and sulfur-containing groups. X may be reactive groups, such as the silanol, amino, thiol, halogen, alkoxy, and alkenyl groups, and the hydrogen radical.

Preference is given to linear polysiloxanes of composition $SiR^2{}_nX_m[SiR^2{}_xX_yO]_zSiR^2{}_oX_p$ (where $0 \le n \le 3$; $0 \le m \le 3$; $0 \le x \le 2$; $0 \le y \le 2$; $0 \le o \le 3$; $0 \le p \le 3$; $1 \le z \le 10000$, where n + m = 3; x + y = 2; o + p = 3), R^2

Particular preference is given to polysiloxanes of composition $SiR^2{}_nX_m[SiR^2{}_xX_yO]_zSiR^2{}_oX_p$ (where $0 \le n \le 3$; $0 \le m \le 1$; $0 \le x \le 2$; $0 \le y \le 2$; $0 \le o \le 3$; $0 \le p \le 1$; $1 \le z \le 1000$, where n + m = 3, x + y = 2, o + p = 3), R^2

the optionally pelletized, of modification The unpelletized, ground, and/or unground precipitated silica with one or more of the organosilanes mentioned 20 may take place in mixtures of from 0.5 to 50 parts, 100 parts of precipitated silica, particular from 1 to 15 parts, based on 100 parts of the reaction precipitated silica, where precipitated silica and organosilane may be carried out 25 during preparation of the mixture (in situ) externally via spray application and subsequent heatconditioning of the mixture, via mixing of the organosilane and the silica suspension with subsequent drying and heat-conditioning (e.g. as in DE 34 37 473 30 and DE 196 09 619), or by the process described in DE 196 09 619 or DE-C 40 04 781.

Suitable organosilicon compounds are in principle any of the bifunctional silanes which firstly can bring about coupling to the filler containing silanol groups and secondly can bring about coupling to the polymer. Amounts usually used as the organosilicon compounds are

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from 1 to 10% by weight, based on the total amount of precipitated silica.

Examples of these organosilicon compounds are:

bis(3-triethoxysilylpropyl) tetrasulfane, bis(3-triethoxysilylpropyl) disulfane, vinyltrimethoxysilane, vinyltriethoxysilane, 3-mercaptopropyltrimethoxysilane, 3-mercaptopropyltriethoxysilane, 3-aminopropyltrimethoxysilane, methoxysilane, 3-aminopropyltriethoxysilane. Other organosilicon compounds have been described in WO 99/09036, EP 1 108 231, DE 101 37 809, DE 101 63 945, DE 102 23 658.

In one preferred embodiment of the invention, bis(3-triethoxysilylpropyl) tetrasulfane may be used as silane.

The silica of the invention may be incorporated as reinforcing filler in elastomer mixtures, tires, or vulcanizable rubber mixtures, in amounts of from 5 to 200 parts, based on 100 parts of rubber, in the form of powder, beads, or pellets, either with silane modification or else without silane modification.

For the purposes of the present invention, rubber 25 mixtures and elastomer mixtures are regarded as equivalent.

The silanol groups on the silica surface function in rubber mixtures as possible chemical reaction partners for a coupling reagent. By way of example, this is a 30 silane, bis(3-triethoxysuch as bifunctional silylpropyl) tetrasulfane, which permits the linkage of the silica to the rubber matrix. The maximum number of silanol groups therefore achieves high probability of coupling between silica and the coupling reagent, and 35 thus high probability of linkage of the silica to the rubber matrix, the final result of this being higher reinforcing potential. The Sears value V₂ is a

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dimension allowing description of the number of silanol groups in the silica, while the BET surface area of a silica describes its specific surface area, which has a major effect on the processing behavior of a compounded other properties its on material. and data relating to the However, the vulcanization. absolute number of silanol groups are not themselves for adequate characterization of sufficient precipitated silica, because precipitated silicas with a high surface area generally have a higher absolute 10 number of silanol groups than precipitated silicas with a low surface area. The important factor is therefore the quotient calculated by dividing the Sears value $\ensuremath{\text{V}}_2$ by the BET. In this way it is possible to represent the reinforcement potential generated via 15 groups per unit of specific surface area introduced.

the elastomer mixtures or rubber fillers in The mixtures may comprise not only mixtures comprising exclusively the silicas of the invention, with or 20 without the abovementioned organosilanes but also one relatively high more other fillers with relatively low reinforcing capability.

- The other fillers used may comprise the following 25 materials:
- Carbon blacks: the carbon blacks to be used here are prepared by the flame black process, furnace black process, or gas black process, and have BET surface areas of from 20 to 200 m^2/g , e.g. SAF, ISAF, HSAF, 30 HAF, FEF, or GPF blacks. Where appropriate, the carbon blacks may also contain heteroatoms, e.g. silicon.
- Fine-particle fumed silicas, e.g. prepared by flame hydrolysis of silicon halides. Where appropriate, the 35 silicas may also be mixed oxides with other metal oxides, such as Al oxides, Mg oxides, Ca oxides, Ba oxides, Zn oxides, or titanium oxides.

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- Other commercially available silicas.
- Synthetic silicates, such as aluminum silicate, or alkaline earth metal silicates, such as magnesium silicate or calcium silicate, with BET surface areas of from 20 to 400 m^2/g and with primary particle diameters of from 10 to 400 nm.
- Synthetic or natural aluminum oxides and synthetic or natural aluminum hydroxides.
- Natural silicates, such as kaolin, and other
 naturally occurring silicon dioxide compounds.
 - Glass fiber and glass fiber products (mats, strands), or glass microbeads.
 - Starch and modified grades of starch.
- Naturally occurring fillers, e.g. clays and siliceous chalk.

Here, as with the amounts used of the organosilanes, the mixing ratio depends on the property profile to be achieved in the finished rubber mixture. A conceivable ratio, also implemented here, between the silicas of the invention and the other abovementioned fillers (including a mixture) is 5-95%.

In one particularly preferred embodiment, from 10 to 150 parts by weight of silicas, composed entirely or to some extent of the silica of the invention, may be used to prepare the mixtures, where appropriate together with from 0 to 100 parts by weight of carbon black, use may also be made of from 1 to 10 parts by weight of an organosilicon compound, based in each case on 100 parts by weight of rubber.

Another important constituent of the rubber mixture, besides the silicas of the invention, the organosilanes, and other fillers, is the elastomers. Mention may be made here of elastomers, natural or synthetic, oil-extended or non-oil-extended, in the form of single polymer or blend with other rubbers,

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examples being natural rubbers, polybutadiene (BR), polyisoprene (IR), styrene/butadiene copolymers having styrene contents of from 1 to 60% by weight, preferably from 2 to 50% by weight (SBR), in particular prepared by means of the solution polymerization process, butyl rubbers, isobutylene-isoprene copolymers (IIR), butadiene-acrylonitrile copolymers having acrylonitrile contents of from 5 to 60% by weight, preferably from 10 to 50% by weight (NBR), partially or fully hydrogenated NBR rubber (HNBR), ethylene-propylene-diene copolymers (EPDM), or else a mixture of these rubbers.

The following additional rubbers may also be used for rubber mixtures with the rubbers mentioned:

15 carboxy rubbers, epoxy rubbers, trans-polypentenamers, halogenated butyl rubbers, 2-chlorobutadiene rubbers, ethylene-vinyl acetate copolymers, ethylene-propylene copolymers, and, where appropriate, also chemically derivatized natural rubber, and also modified natural rubbers.

By way of example, W. Hofmann, "Kautschuktechnologie" [Rubber technology], Genter Verlag, Stuttgart, 1980, describes preferred synthetic rubbers.

Anionically polymerized SSBR rubbers (solution SBRs) with a glass transition temperature above -50°C are of particular interest for producing the tires of the invention, as are mixtures of these with diene rubbers.

The silicas of the invention, with or without silane,
30 may be used in any of the applications of rubbers, for
example moldings, tires, tire treads, conveyor belts,
gaskets, drive belts, hoses, shoe soles, cable
sheathing, roller coverings, damping elements, etc.

35 The incorporation of this silica and the preparation of the mixtures comprising this silica, take place in an internal mixer or on a roll mill, preferably at 80-200°C, in the manner conventional in the rubber

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industry. The form in which the silicas are used may be either powder, beads, or pellets. Again, there is no difference between the silicas of the invention and the known pale-color fillers.

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The rubber vulcanizates of the invention may comprise the usual amounts of other rubber auxiliaries, such as reaction accelerators, antioxidants, heat stabilizers, light stabilizers, ozone stabilizers, processing aids, plasticizers, tackifiers, blowing agents, dyes, pigments, waxes, extenders, organic acids, retarders, metal oxides, and also activators, such as triethanolamine, polyethylene glycol, and hexanetriol. These compounds are known in the rubber industry.

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The amounts which may be used of the rubber auxiliaries are those which are known, and depend, inter alia, on the intended use. Examples of usual amounts are from 0.1 to 50% by weight, based on the rubber used. Sulfur or sulfur-donor substances may be used as crosslinkers. 20 The rubber mixtures of the invention may moreover Examples comprise vulcanization accelerators. accelerators are suitable primary mercaptobenzothiazoles, sulfenamides, thiurames, and dithiocarbamates in amounts of from 0.5 to 3 왕 25 weight. Examples of coaccelerators are guanidines, thioureas, and thiocarbonates in amounts of from 0.5 to 5% by weight. The amounts of sulfur which may usually be used are from 0.1 to 10% by weight, preferably from 1 to 3% by weight, based on the rubber used. 30

The silicas of the invention may be used in rubbers which are crosslinkable using accelerators and/or sulfur, or else peroxidically crosslinkable.

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The rubber mixtures of the invention may be vulcanized at temperatures of from 100 to 200°C, preferably from 130 to 180°C, where appropriate under a pressure of

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from 10 to 200 bar. Known mixing assemblies, such as roll mills, internal mixers, and mixing extruders, may be used for blending of the rubbers with the filler, where appropriate with rubber auxiliaries, and with the organosilicon compound.

The rubber mixtures of the invention are suitable for producing moldings, for example for producing pneumatic tires, tire treads for tires for summer, winter, or year-round use, car tires, tires for utility vehicles, motorcycle tires, tire carcass components, cable sheathing, hoses, drive belts, conveyor belts, roller coverings, shoe soles, gasket ring seals, and damping elements.

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An advantage of the silicas of the invention is that they are highly dispersible and that their low microporosity gives them higher rubber activity with respect to the rubber vulcanizates, and that they therefore provide greater reinforcement to the rubber vulcanizates.

When comparison is made with an identical rubber mixture using a prior-art silica, the silicas of the invention also exhibit advantages in vulcanization time.

The rubber mixtures of the invention are particularly suitable for producing tire treads for car tires and for motor cycle tires, and also for producing utility vehicle tires with reduced rolling resistance together with good abrasion resistance and good suitability for winter conditions.

35 The rubber mixtures of the invention are also suitable, without addition of organosilicon compounds, blended with a typical tire tread carbon black, for improving the cut & chip performance of tires on construction

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machinery, on agricultural machinery, or tires used in mining. (For definitions and further details see "New insights into the tear mechanism" and references cited therein, presented by Dr. W. Niedermeier at Tire Tech 2003 in Hamburg.)

The reaction conditions and the physico-chemical data for the precipitated silicas of the invention are determined using the following methods:

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Determination of moisture level of silicas

Based on ISO 787-2, this method is used to determine the volatile content (hereinafter termed moisture level for simplicity) of silica after drying for 2 hours at 105°C. This loss on drying is generally composed mainly of moisture.

Method

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10 g of the silica in powder, bead, or pellet form are weighed out precisely to 0.1 mg into a dry glass weighing vessel with ground-glass lid (diameter 8 cm, height 3 cm) (specimen weight E). With the lid open, the specimen is dried for 2 h at $105 \pm 2^{\circ}$ C in a drying

- the specimen is dried for 2 h at 105 ± 2°C in a drying cabinet. The glass weighing vessel is then sealed, and cooled to room temperature in a desiccator cabinet, using silica gel as drying agent. The specimen is weighed to give the final weight A.
- 30 The moisture level in % is determined as ((E in g A in g)*100%)/(E in g).

Determination of modified Sears value of silicas

35 The modified Sears value (hereinafter termed Sears value V_2) can be determined by titrating silica with potassium hydroxide solution in the range from pH 6 to pH 9, and is a measure of the number of free hydroxy

groups.

The following chemical reactions underlie the determination method, "Si"-OH being intended to symbolize a silanol group of the silica:

"Si"-OH + NaCl \rightarrow "Si"-ONa + HCl HCl + KOH \rightarrow KCl + H₂O.

corresponds to V_1' .

Method

10.00 g of a silica in powder, bead or pellet form, 10 having a moisture level of 5 ± 1%, are comminuted for 60 seconds, using an M 20 IKA universal mill (550 W; 20 000 rpm). Where appropriate, the moisture content of the starting substance has to be adjusted by drying at 105°C in the drying cabinet or uniform moistening, with 15 repeated comminution. 2.50 g of the silica thus treated are weighed out at room temperature into a titration vessel and mixed with 60.0 ml of analytical grade methanol. After complete wetting of the specimen, 40.0 ml of deionized water are added, and the mixture 20 is dispersed, using a T 25 UltraTurrax mixer (KV-18G mixing shaft, 18 mm diameter) for 30 seconds at a rotation rate of 18 000 rpm. The particles of specimen adhering to the edge of the vessel and to the stirrer are flushed into the suspension, using 100 25 deionized water, and a water bath with thermostat is used to control the temperature of the mixture to 25°C. tester (Knick 766 Calimatic pH meter with The pH temperature sensor) and pH electrode (Schott N7680 combination electrode) are calibrated using buffer 30 solutions (pH 7.00 and 9.00) at room temperature. Using the pH meter, the initial pH of the suspension is first measured at 25°C, and then, depending on the result, mol/1)solution (0.1 potassium hydroxide hydrochloric acid solution (0.1 mol/l) is used to 35 adjust the pH to 6.00. The consumption of KOH solution HCl solution in ml required to reach pH 6.00

20.0 ml of sodium chloride solution (250.00 g of analytical grade NaCl made up to 1 l using deionized water) are then metered in. The titration is then continued as far as pH 9.00, using 0.1 mol/l KOH. The consumption of KOH solution in ml required to reach pH 9.00 corresponds to $V_2{}^\prime$.

The volumes V_1 ' and V_2 ' are first then normalized for a theoretical specimen weight of 1 g, and are then multiplied by 5, thus giving V_1 and the Sears value V_2 in the units ml/(5 g).

Determination of BET surface area

The specific nitrogen surface area (hereinafter termed 15 BET surface area) of the silica in powder, bead, or pellet form is determined to ISO 5794-1/Annex D, using an AREA-meter (Ströhlein, JUWE).

CTAB surface area determination

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The method is based on the adsorption of CTAB (Nhexadecyl-N, N, N-trimethylammonium bromide) the on "external" surface of the silica, also termed the "rubber-active surface", and is based on ASTM3765 or NFT 45-007 (Section 5.12.1.3). CTAB is adsorbed aqueous solution, with stirring and treatment with ultrasound. Excess, unadsorbed CTAB is determined by back-titration with NDSS (solution of the sodium salt of dioctyl sulfosuccinate, "Aerosol OT" solution), using a titroprocessor, the end point being given by maximum clouding of the solution, and determined using a phototrode. The temperature during all of operations carried out is 23-25°C, in order to prevent CTAB from crystallizing out. The following equation underlies the back-titration reaction:

 $(C_{20}H_{37}O_4) SO_3Na + BrN(CH_2)_3 (C_{16}H_{22}) \rightarrow (C_{20}H_{37}O_4) SO_3N(CH_3)_3 (C_{16}H_{33}) + NaBr \\ NDSS \\ CTAB$

Equipment

METTLER Toledo DL 55 titroprocessor and METTLER Toledo DL 70 titroprocessor, in each case equipped with: DG 111 pH electrode produced by Mettler, and DP 550 phototrode produced by Mettler 100 ml polypropylene titration beaker 150 ml glass titration vessel with lid Pressure-filtration equipment, capacity 100 ml Membrane filter composed of cellulose nitrate, pore size 0.1 μ m, diameter 47 mm, e.g. Whatman (order no.

15 Reagents

7181-004)

The solutions of CTAB (0.015 mol/l in deionized water) and NDSS (0.00423 mol/l in deionized water) are purchased in ready-to-use form (Kraft, Duisburg: order no. 6056.4700 CTAB solution, 0.015 mol/l; order no. 6057.4700 NDSS solution, 0.00423 mol/l), stored at 25°C, and used within one month.

Method

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Blind titration

The consumption of NDSS solution for the titration of 5 ml of CTAB solution is to be tested once daily prior to each series of tests. For this, the phototrode is adjusted to 1000 ± 20 mV (corresponding to 100% transparency) prior to starting the titration.

Exactly 5.00 ml of CTAB solution are pipetted into a titration beaker, and 50.0 ml of deionized water are added. The titration with NDSS solution uses the test method familiar to the person skilled in the art, with stirring, using the DL 55 titroprocessor until max. clouding of the solution has been reached. The

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consumption V_{I} of NDSS solution is determined in ml. Three determinations are to be carried out for each titration.

5 Adsorption

10.0 g of the silica in powder, bead or pellet form, with a moisture content of 5 \pm 2% (the moisture content being adjusted, where appropriate, by drying at 105°C in the drying cabinet or uniform moistening), 10 comminuted for 30 seconds, using a mill (Krups KM 75, item no. 2030-70). Exactly 500.0 mg of the comminuted specimen are transferred into a 150 ml titration vessel with magnetic stirrer bar, and exactly 100.0 ml of CTAB solution are metered in. The titration vessel is sealed 15 with a lid, and the mixture is stirred for 15 minutes, using a magnetic stirrer. Hydrophobic silicas stirred, using a T 25 UltraTurrax mixer (KV-18G mixer shaft, 18 mm diameter), at 18 000 rpm for a maximum of 1 min to give complete wetting. The titration vessel is 20 screw-fastened to the DL 70 titroprocessor, and the pH of the suspension is adjusted to a value of 9 + 0.05, using KOH (0.1 mol/l). The suspension is exposed for 4 minutes to ultrasound, in the titration vessel, in an 106 S from Bandelin, ultrasound bath (Sonorex RK 25 immediately 35 KHz) at 25°C. This is followed filtration under pressure through a membrane filter, the nitrogen pressure being 1.2 bar. The first 5 ml of filtrate are discarded.

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Titration

5.00~ml of the remaining filtrate are pipetted into a 100~ml titration beaker and made up to 50.00~ml, using deionized water. The titration beaker is screw-fastened to the DL 55~titroprocessor, and the mixture is titrated with NDSS solution, with stirring, until maximum clouding has been reached. The consumption V_{II}

of NDSS solution in ml is determined. Each clouding point is to be determined three times.

Calculation

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The measured values

 $V_{\text{I}} =$ consumption of NDSS solution in ml when the blind specimen is titrated

 V_{II} consumption of NDSS solution in ml when the filtrate is used

give:

 $V_{\rm I}/V_{\rm II}$ = amount of CTAB in the blind specimen/remaining amount of CTAB present in the filtrate specimen.

The amount N of adsorbed CTAB in g is therefore:

 $N = ((V_I - V_{II}) * 5.5 g * 5 ml)/V_I * 1000 ml).$

Because only 5 ml were titrated from 100 ml of 10 filtrate, and 0.5 kg of silica of a defined moisture level was used, and the area required by 1 g of CTAB is $578435 \times 10^{-3} \text{ m}^2$ and:

CTAB surface area (not corrected for water) in $m^2/g = (N^*20^*578.435 m^2/g)/(0.5 g)$, and

15 CTAB surface area (not corrected for water) in $m^2/g = ((V_I-V_{II})^* 636.2785 m^2/g)/V_I$.

The CTAB surface area is based on the anhydrous silica, and the following correction is therefore made:

CTAB surface area in $m^2/g = (CTAB \text{ surface area (not corrected for water) in } m^2/g * 100%)/(100% - moisture level in %).$

Determination of DBP absorption

DBP absorption (DBP number) is determined as follows by a method based on the standard DIN 53601, and is a measure of the absorbency of the precipitated silica:

Method

12.50 g of silica in powder or bead form with 0-10% moisture content (the moisture content being adjusted, where appropriate, by drying at 105°C in the drying cabinet) are placed in the kneader chamber (item no. 279061) of the Brabender "E" Absorptometer (with no damping of the output filter of the torque sensor). In the case of pellets, the sieve fraction from 3.15 to 10 1 mm (Retsch stainless-steel sieve) is used (by using slight pressure from a plastics spatula to press the pellets through the 3.15 mm-pore-width sieve). While the material is constantly mixed (rotation rate of kneader paddles: 125 rpm), dibutyl phthalate is added 15 dropwise at a rate of 4 ml/min at room temperature into the mixture by way of the "Brabender T 90/50 Dosimat". Incorporation by mixing is achieved with only low power consumption, and is followed by means of the digital display. Toward the end of the determination, the 20 mixture becomes pasty, and this is indicated by a steep rise in the power consumption. When the display shows 600 digits (0.6 Nm torque), both the kneader and the DBP feed are switched off via an electrical contactor. The synchronous motor for the DBP supply has coupling 25 to a digital meter, thus permitting read-off of DBP consumption in ml.

Evaluation

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DBP absorption is stated in g/(100~g), and is calculated from the measured DBP consumption by using the following formula. The density of DBP at 20° C is typically 1.047~g/ml.

35 DBP absorption in g/(100 g) = ((DBP consumption in ml)* (DBP density in g/ml) * 100)/(12.5 g).

Defined DBP absorption relates to the anhydrous, dried

silica. If moist precipitated silicas are used, the value is to be corrected by means of the following correction table. The correction value, corresponding to the water content, is added to the DBP value determined experimentally; by way of example, an addition of $33 \, \text{g/(100 g)}$ for DBP absorption would be implied if the water content were 5.8%.

Correction table for dibutyl phthalate absorption - 10 anhydrous -

	.% water				
% water	. 0	. 2	. 4	. 6	. 8
0	0	2	4	5	7
1	9	10	12	13	15
2	16	18	19	20	22
3	23	24	26	27	28
4	28	29	29	30	31
5	31	32	32	33	33
6	34	34	35	35	36
7	36	37	38	38	39
8	39	40	40	41	41
9	42	43	43	44	44
10	45	45	46	46	47

pH determination

The method, based on DIN EN ISO 787-9, serves for determination of the pH of an aqueous suspension of silicas at 20°C. To this end, an aqueous suspension is prepared from the specimen to be studied. After brief shaking of the suspension, its pH is determined by means of a previously calibrated pH meter.

Method

Prior to making the pH measurement, daily calibration of the pH tester (Knick 766 Calimatic pH meter with

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temperature sensor) and of the pH electrode (Schott N7680 combination electrode, is required, using the buffer solutions, at 20°C. The calibration function is to be selected in such a way that the two buffer solutions used include the expected pH of the specimen (buffer solutions having pH 4.00 and 7.00, pH 7.00 and 9.00, and, where appropriate, pH 7.00 and 12.00). If pellets are used, 20.0 g of silica are first comminuted for 20 s by means of a mill (Krups KM 75, item No. 2030-70).

5.00 g of silica in powder or bead form, with $5 \pm 1\%$ moisture content (where appropriate, the moisture content being adjusted by drying at 105°C in the drying cabinet or by uniform wetting prior to any comminution) are precisely weighed out to 0.01 g on a precision balance into a previously tared wide-necked glass flask. 95.0 ml of deionized water are added to the specimen. An automatic shaker (Gerhardt, LS10, 55 W, level 7) is then used to shake the suspension in the sealed vessel for a period of 5 minutes at temperature. The pH is measured directly after shaking. this end, the electrode is rinsed first then with a portion of the deionized water and suspension, and then dipped into the suspension. A magnetic stirrer bar is then added to the suspension, and the pH measurement is made at a constant stirring rate with formation of slight vortex in the suspension. When the pH meter displays a constant value, the pH is read off from the display.

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The method when hydrophobic silica is used is similar, but in that case 5.00 g of the specimen, after comminution where appropriate and with $5 \pm 1\%$ moisture content, are weighed out precisely to 0.01 g on the precision balance into a previously tared wide-necked glass flask. 50.0 ml of analytical grade methanol and 50.0 ml of deionized water are added, and an automatic shaker (Gerhardt, LS10, 55 W, level 7) is then used to

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shake the suspension in the sealed vessel for a period of 5 minutes at room temperature. The pH is likewise measured with stirring, but after precisely 5 min.

5 Determination of solids content of filter cake

This method is used to determine the solids content of filter cake via removal of the volatile content at $105\,^{\circ}\text{C}$.

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Method

100.00 g of the filter cake are weighed out into a dry, tared porcelain dish (diameter 20 cm) (specimen Where appropriate, the filter cake weight E). 15 in order to comminuted, using a spatula, separate fragments of not more than 1 $\ensuremath{\text{cm}^3}$. The specimen is dried to constant weight at 105 ± 2°C in a drying specimen is then cooled to cabinet. The temperature in a desiccator cabinet, using silica gel 20 as drying agent. The material is weighed to determine the final weight A.

The solids content in % is determined as 100% - (((E in g - A in g)*100%)/(E in g)).

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Determination of electrical conductivity

The electrical conductivity (C) of silicas is determined in aqueous suspension.

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Method

If pellets are used, 20.0 g of silica are first comminuted by means of a mill (Krups, KM 75, item No. 2030-70) for 20 s. 4.00 g of silica in powder or bead form, with 5 ± 1% moisture content (the moisture content being adjusted, where appropriate, by drying at 105°C in the drying cabinet or uniform wetting prior to

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any comminution) are suspended in 50.0 ml of deionized water, and heated for 1 min to 100°C. The specimen cooled to 20°C is made to precisely 100 ml and homogenized by rotary shaking.

The measurement cell of the LF 530 (WTW) conductivity tester is flushed with a small amount of specimen before the LTA01 measurement cell is dipped into the suspension. The value displayed corresponds to the conductivity at 20°C, because the TFK 530 external temperature sensor automatically compensates for the temperature. Both this temperature coefficient and the cell constant k are to be checked prior to each series of tests.

The calibration solution used comprises 0.01 mol/l potassium chloride solution (C at 20°C = 1278 μ S/cm).

Determination of solids content of precipitation suspension

20 The solids content of the precipitation suspension is determined gravimetrically after filtration of the specimen.

Method

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100.0 ml of the homogenized precipitation suspension $(V_{\text{suspension}})$ are measured out at room temperature with aid of a measuring cylinder. The specimen filtered off with suction by way of a round filter (Schleicher & Schuell 572) in a porcelain suction 30 filter funnel, but not sucked dry, so as to avoid cracking of the filter cake. The filter cake is then washed with 100.0 ml of deionized water. The suctionfilter cake is of the leached filtration completed, and the filter cake is transferred into a 35 tared porcelain dish and dried to constant weight at 105 \pm 2°C in a drying cabinet. The weight of the dried silica (m_{specimen}) is determined.

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The solids content is determined as: solids content in $g/l = (m_{specimen} g)/(V_{suspension} in l)$.

Determination of alkali value

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The alkali value (AV) determined is the consumption in ml of hydrochloric acid (using a specimen volume of 50 ml, 50 ml of dist. water, and a hydrochloric acid concentration of 0.5 mol/l) during direct potentiometric titration of alkaline solutions or alkaline suspensions as far as pH 8.30. The result reveals the free alkali content of the solution or suspension.

15 Method

The pH device (Knick, Calimatic 766 pH meter with temperature sensor) and the pH electrode (Schott N7680 electrode) are calibrated at combination temperature with the aid of two buffer solutions (pH = 20 7.00 and pH = 10.00). The combination electrode is dipped into the test solution or test suspension, the temperature of which has been controlled to 40°C, and which is composed of 50.0 ml of specimen and 50.0 ml of deionized water. Hydrochloric acid solution whose 25 concentration is 0.5 mol/l is then added dropwise until constant pH of 8.30 is obtained. Because the equilibrium between the silica and the free alkali content is established only slowly, a waiting time of 15 min is required before final read-off of the acid 30 For the amounts and concentrations consumption. selected, the read-off of hydrochloric acid consumption in ml directly corresponds to the alkali value, which is a dimensionless quantity.

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The following examples are intended to provide further illustration of the invention but not to limit its scope.

Example 1

Silica I

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1550 l of water and 141.4 kg of water glass (density) 1.348 kg/l, 27.0% SiO_2 , 8.05% Na_2O) form an initial charge in a reactor made from stainless steel with a propeller-stirrer system and jacket heating. 5.505 kg/min of the abovementioned water glass and about 0.65 kg/min of sulfuric acid (density 1.83 kg/l, 96% $\mathrm{H}_{2}\mathrm{SO}_{4})$ are then metered in with vigorous stirring at 92°C over a period of 80 minutes. This metering of sulfuric acid is regulated in such a way that the alkali value prevailing in the reaction mixture is 20. The water glass addition is then stopped, and the addition of sulfuric acid is continued until a pH of 9.0 (measured at room temperature) has been achieved. The sulfuric acid addition is stopped, and stirring of the resultant suspension is continued for 60 minutes at 90°C. Addition of sulfuric acid is then restarted, and a final pH of 3.5 (measured at room temperature) is established.

The resultant suspension is filtered, using a membrane filter press, and the product is washed with water. The filter cake, with 23% solids content, is liquidized, using aqueous sulfuric acid and a shearing assembly. The silica feed with 19% solids content and with a pH of 3.7 is then spray-dried with addition of ammonia.

30 The resultant powder Ia has a BET surface area of $114~\text{m}^2/\text{g}$ and a CTAB surface area of $123~\text{m}^2/\text{g}$. The spraytower-dried microbead product Ib obtained as an alternative has a BET surface area of $126~\text{m}^2/\text{g}$ and a CTAB surface area of $123~\text{m}^2/\text{g}$.

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Silica II

1550 l of water and 141.4 kg of water glass (density

1.348 kg/l, 27.0% SiO₂, 8.05% Na₂O) form an initial charge in a reactor made from stainless steel with a propeller-stirrer system and jacket heating. 5.505 kg/min of the abovementioned water glass and about 0.65 kg/min of sulfuric acid (density 1.83 kg/l, 96% H₂SO₄) are then metered in with vigorous stirring at 92°C over a period of 80 minutes. This metering of sulfuric acid is regulated in such a way that the alkali value prevailing in the reaction mixture is 20.

The water glass addition is then stopped, and the addition of sulfuric acid is continued until a pH of 7.0 (measured at room temperature) has been achieved. The sulfuric acid addition is stopped, and stirring of the resultant suspension is continued for 60 minutes at

90°C. Addition of sulfuric acid is then restarted, and a final pH of 3.5 (measured at room temperature) is established.

The resultant suspension is filtered, using a membrane filter press, and the product is washed with water. The 20 filter cake, with 22% solids content, is liquidized, using aqueous sulfuric acid and a shearing assembly. The silica feed with 19% solids content and with a pH of 3.6 is then spray-tower-dried with addition of ammonia.

The resultant microbead product has a BET surface area of 116 $\rm m^2/g$ and a CTAB surface area of 117 $\rm m^2/g$.

Silica III

1.348 kg/l, 27.0% SiO₂, 8.05% Na₂O) form an initial charge in a reactor made from stainless steel with a propeller-stirrer system and jacket heating. 5.505 kg/min of the abovementioned water glass and about 0.65 kg/min of sulfuric acid (density 1.83 kg/l, 96% H₂SO₄) are then metered in with vigorous stirring at 92°C over a period of 100 minutes. This metering of sulfuric acid is regulated in such a way that the

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alkali value prevailing in the reaction mixture is 20. The water glass addition is then stopped, and the addition of sulfuric acid is continued until a pH of 9.0 (measured at room temperature) has been achieved. The sulfuric acid addition is stopped, and stirring of the resultant suspension is continued for 60 minutes at 90°C. Addition of sulfuric acid is then restarted, and a final pH of 3.5 (measured at room temperature) is established.

The resultant suspension is filtered, using a membrane filter press, and the product is washed with water. The filter cake, with 23% solids content, is liquidized, using aqueous sulfuric acid and a shearing assembly. The silica feed with 19% solids content and with a pH of 3.8 is then spray-dried with addition of ammonia. The resultant product has a BET surface area of 102 m²/g and a CTAB surface area of 104 m²/g.

The physical data for the above silicas are given in 20 the table below:

		Silica	Silica	Silica	Silica	REF
		Ia	Ib	II	III	Ultrasil
						7000 GR
Finishing		PH 9	9 Hq	рН 7	рн 9	
Moisture level	[%]	4.5	5.1	5.1	5.0	5.4
рН	[-]	5.6	5.5	5.5	5.6	6.2
Conductivity	[µS/cm]	610	550	600	590	620
BET	[m ² /g]	114	126	116	102	174
CTAB	[m ² /g]	123	123	117	104	163
BET/CTAB	[-]	0.93	1.02	0.99	0.98	1.07
DBP	[g/(100 g)]	278	258	274	256	228
Sears value V ₂	[ml/(5 g)]	22	21	22	22	16
Sears value	$[m1/(5 m^2)]$	0.193	0.167	0.190	0.216	0.092
V ₂ /BET					ĺ	

Example 2

The precipitated silicas Ib and II of the invention from Example 1 were studied in an SSBR/BR rubber mixture. The readily dispersible silica Ultrasil 7000

GR (Reference) from Degussa AG was selected as prior art.

The mixing specification used for the rubber mixtures is given in Table 1 below. The unit phr here means parts by weight, based on 100 parts of the untreated rubber used.

Table 1

Table 1	
1st stage	
Buna VSL 5025-1	96
Buna CB 24	30
Silica	80
x50-S	12.8
ZnO	2
Stearic acid	2
Naftolen ZD	10
Vulkanox 4020	1.5
Protector G 3108	1
2nd stage	
Stage 1 batch	
3rd stage	
Stage 2 batch	
Vulkacit D/C	2.0
Vulkacit CZ/EG	1.5
Perazit TBZTD	0.2
Sulfur	1.2

The polymer VSL 5025-1 is a solution-polymerized SBR copolymer from Bayer AG with a styrene content of 25% by weight and a butadiene content of 75% by weight. The copolymer comprises 37.5 phr of oil and has a Mooney viscosity (ML 1+4/100°C) of 50 ± 4. The polymer Buna CB 24 is a cis-1,4-polybutadiene (neodymium grade) from Bayer AG with cis-1,4 content of at least 97% and a Mooney viscosity of 44 ± 5.

X50-S is a 50/50 blend of Si 69 [bis(3-triethoxysilyl-propyl) tetrasulfide] and carbon black obtainable from

20 Degussa AG.

The aromatic oil used comprises Naftolen ZD from Chemetall. Vulkanox 4020 is 6PPD from Bayer AG and Protektor G 3108 is an ozone-protection wax from HB-Fuller GmbH. Vulkacit D/C (DPG) and Vulkacit CZ/EG (CBS) are commercially available products from Bayer AG. Perkazit TBZTD is obtainable from Akzo Chemie GmbH. The rubber mixtures are prepared in an internal mixer using the mixing instructions in Table 2. Table 3 gives the methods used for rubber testing. The mixtures are vulcanized at 165°C for 20 minutes. Table 4 shows the results of testing on the vulcanized rubber.

Table 2

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	Stage 1				
Settings	Settings				
Mixing assembly	Werner & Pfleiderer 1.5N				
Rotation rate	70 rpm				
Friction	1:1.11				
Ram pressure	5.5 bar				
Capacity	1.6 l				
Fill level	0.73				
Chamber temp.	70°C				
Mixing procedure					
0-0.5 min	Buna VSL 5025-1 + Buna CB 24				
0.5-1.5 min	1/3 filler, X50-S				
1.5-2.5 min	1/3 filler				
2.5-3.5 min	1/3 filler, remaining stage 1 constituents				
3.5-4.5 min	mixing				
4.5-6.0 min	mixing and discharge				
Storage	2: h at room temperature				

Stage 2				
Settings				
Mixing assembly	as in stage 1 except:			
Rotation rate	80 rpm			
Chamber temp.	80°C			
Fill level	0.71			
Mixing procedure				
0-5 min	-5 min stage 1 batch, maintain batch temperature			
	140-150°C by varying rotation rate			
5 min	discharge			
Storage	4 h at room temperature			

Stage 3			
Settings	•		
Mixing assembly	as in stage 1 except:		
Rotation rate	40 rpm		
Chamber temp.	50°C		
Fill level	0.69		
Mixing procedure			
0-2 min	stage 2 batch, stage 3 constituents		
2 min	discharge and form milled sheet on		
	laboratory roll mill,		
	(diameter 200 mm, length 450 mm, roll		
	temperature 50°C)		
	homogenization:		
cut the material 3 times toward the left			
3 times toward the right			
fold the material over 5 times with narro			
	roll gap (1 mm) and		
	5 times with wide roll gap (3.5 mm) and		
	peel milled sheet away		

Table 3

Standard/conditions Physical testing DIN 53529/3, ISO 6502 Vulcameter testing, 165°C Torque difference Dmax-Dmin [dNm] t10% and t90% [min] DIN 53504, ISO 37 Ring tensile test, 23°C Stress values 100% and 300% [Mpa] Reinforcement factor: stress value 300%/100%[-] DIN 53 505 Shore A hardness, 23°C[-] Ball rebound [%], 0°C and 60°C DIN EN ISO 8307 steel ball, 19 mm, 28 g see text Dispersion coefficient [%] DIN 53 513, ISO 2856 Viscoelastic properties, 0 and 60°C, 16 Hz, initial force 50 N and amplitude force 25 N Test value recorded after 2 min. of test time, i.e. 2 min. of conditioning Complex modulus E* [Mpa] Loss factor tan δ [-]

The dispersion coefficient was determined using the inc. Medalia correction surface topography 5 (A. Wehmeier, "Filler Dispersion Analysis by Topography Measurements" Technical Report TR 820, Degussa AG, Advanced Fillers and Pigments Division). The dispersion coefficient thus determined correlates at reliability level > 0.95 with the optically determined dispersion 10 coefficient, for example as determined by the Deutschen Kautschuktechnologie e.V. [Germany für Institut Institute for Rubber Technology], Hanover, Germany (H. Geisler, Bestimmung der Mischgüte, presented at DIK Workshop, 27-28 November 1997, Hanover, Germany). 15

Table 4

Data for untreated mixture	REF	Silica Ib	Silica II
Dmax-Dmin	16.4	16.5	16.1
t10%	1.4	1.8	1.8
t90%	6.2	5.5	5.6
Vulcanizate data			·
100% stress	1.7	2.1	2.1
300% stress	9.2	12.3	12.4
300%/100% stress	5.4	5.9	5.9
Shore A hardness	62	64	63
Ball rebound 0°C	15.8	15.1	14.8
Ball rebound 60°C	62.5	64.2	64.4
E* (0°C)	19.2	19.0	18.2
E* (60°C)	7.7	7.9	7.7
tan δ (0°C)	0.475	0.460	0.454
tan δ (60°C)	0.124	0.108	0.106
Dispersion coefficient	94	96	97

As can be seen from the data in Table 4, the Silica Ib and Silica II mixtures with the silicas of the invention have a lower vulcanization time t90% than the reference mixture REF, due to lower microporosity and therefore less surface-adsorption of accelerator. The advantageous higher activity of these silicas reflected in the higher 100% and 300% stress values and 10 increased reinforcement factor. The dynamic properties of the mixtures with the silicas of the invention have likewise been improved. Ball rebound 0°C is found to be lower, implying improved wet skid resistance, and ball rebound 60°C is found to be 15 higher, indicating better rolling resistance of a tire tread. The dynamic stiffnesses at 0 and 60°C are at the same level as those of the reference mixture, while the hysteresis loss tan δ (60°C) has an advantageously been lowered, indicating lower rolling resistance. In 20 addition, the quality of dispersion of the silicas of the invention is extraordinarily high and, inter alia,

better than that of the silica Ultrasil 7000 GR of the prior art, which itself has good dispersibility, thus achieving advantages in road abrasion.